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The ultimate goal of this program is to discover selective and potent drugs targeted to prevent or relieve the neurotoxic actions of botulinum neurotoxin (BoNT) A. A major goal of this program is the identification of open channel blockers as a single class of drugs that would be effective against all BoNT isoforms. The major focus thus far has been the implementation of a reliable and robust high-throughput screen for blockers specific for BoNT. This facet of the program involves the use of the VIPR™ -Voltage/Ion Probe Reader, a proven strategy for high-throughput screening, using nerve growth factor-differentiated pheochromocytoma PC12 cells in which BoNTA forms channels with similar properties to those previously characterized in lipid bilayers. The fidelity of the assay relies on fluorescence measurements of membrane potential changes as an index of open BoNT channels and increased cation conductance. The immediate task is to select mixtures from synthetic combinatorial libraries with high blocking activity to deconvolute and identify the most potent compounds. We consider the BoNT channel as a validated target for intervention aimed to inhibit the translocation of the light chain into the cytosol and therefore to attenuate the BoNT neurotoxicity.

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INTRODUCTION

This program examines innovative approaches and powerful new technologies to identify selective and potent agents directed to prevent or relieve the neuroparalytic toxic actions of botulinum toxin A (BoNTA)¹. The focus is on the ion channel forming activity of BoNTs as a validated target to screen for inhibitors of the translocation of the light chain into the cytosol and therefore to attenuate the BoNT neurotoxicity. The key of the program is based on our discovery that the heavy chain (HC) of BoNT acts as both a channel and a transmembrane chaperone for the light chain (LC) to ensure a translocation competent conformation during its transit from the acidic endosome into the cytosol - its site of action (Korazova, L. & Montal, M. *Nature Struct Biol* **10**, 13-18, 2003. *Appendix #1*)². This is an exciting time to focus on innovative technologies to uncover lead compounds that may represent a potential new generation of useful and safe antidotes for BoNTs.

Two major aims, representing different technologies, are pursued:

- Task 1: Use of synthetic combinatorial libraries (SCL)³ to discover specific inhibitors of the channel activity of BoNTs.
- Task 2: Use of the VIPRTM -Voltage/Ion Probe Reader⁴, for high-throughput screening (HTS) for open channel blockers of BoNTA channel on pheochromocytoma PC12 cells.

The combination of these two strategies converging on the channel activity of BoNT will optimize the likelihood of selecting individual compounds from the libraries in a practical time encompassing the duration of the grant. Evidently, the two tasks are intricately connected. However, inhibitor discovery requires the implementation of a robust and reliable HTS. This has been the main focus of the program during the first year and is the subject matter of this report.

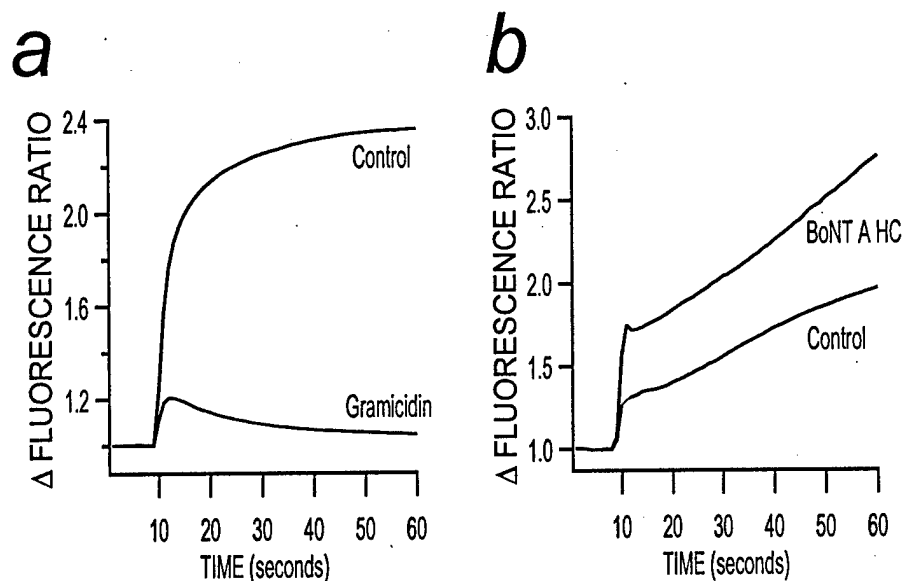
BODY

Our focus has been the application of Fluorescence Resonance Energy Transfer (FRET) assays on nerve growth factor (NGF)-treated PC12 cells⁵ to screen for blockers of BoNT HC channel activity. The HTS assay involves a Vertex, formerly known as Aurora Biosciences, proprietary phospholipid linked coumarin dye (CC2DMPE) paired with voltage sensing oxonol dyes, in addition to custom detection machinery, the VIPRTM-Voltage/Ion Probe Reader⁴, as described in the application. The basic concept underlying this FRET assay is as follows: a coumarin-linked phospholipid-CC2DMPE, when added to cells, adsorbs to the outer lipid monolayer of the cell membrane bilayer and serves as a FRET donor to a mobile, lipid soluble oxonol dye, a bis-(1,3-dialkylthiobarbituric acid) trimethine oxonol (DisBAC2(3), the FRET acceptor. Excitation of the donor results in FRET provided the oxonol dye is in proximity to the donor. When cells are depolarized, the anionic oxonol dye responds to the transmembrane potential by diffusing away toward the inner monolayer of the bilayer, thereby increasing its separation from the membrane anchored donor and leading to a decrement in FRET.

The ratio change (donor/acceptor) is reversed upon repolarization, with oxonol translocation across the bilayer in the ms time range thereby allowing voltage-sensitive FRET to report fast voltage changes. For this assay, cells are plated into 96 well black walled tissue culture plates coated with 1 % Matrigel (BD Biosciences, growth factor reduced formulation). Using a "Hi K⁺ depolarization" protocol to be described, cell densities of 15 % for 2 day NGF-treated PC12 cells were determined to be optimal at final dye concentrations of 10 μ M for CC2DMPE and 2 μ M for DisBAC2(3). The cells are loaded with CC2DMPE in presence of Pluronic F-127, an inert polymer which accelerates loading of the fatty acid linked coumarin into the cell membrane in an aqueous buffer system. After the coumarin wash, using an automatic cell washer program designed for a 96 well plate, the cells are loaded with oxonol and Pluronic F-127 at pH 7.4. BoNT HC is then added to designated wells at a concentration of at least 5 μ g ml⁻¹ to allow for binding of the protein to the cell outer membrane and incubated 20 min at 22-24 °C in the dark. Next, in order to emulate the conditions prevalent at endosomes, the cells are exposed to an acidic environment for 5 min by adding an equal volume oxonol containing a concentration of citrate which brings the final pH to 5.5. To maximize FRET, cells are returned to pH 7.4 in oxonol immediately prior to the assay on the VIPR.

To calibrate the assay, we first proceeded to use a "Hi K⁺ depolarization" protocol in presence of the well-known channel forming peptide gramicidin A⁶. Accordingly, the cells are loaded with oxonol in a solution that contains (in mM) NaCl 160, KCl 4.5, CaCl₂ 2, MgCl₂ 1, Hepes 10, Glucose 10 (pH 7.4 with NaOH). Depolarization in "Hi K⁺" buffer that contains (in mM) KCl 164.5, CaCl₂ 2, MgCl₂ 1, Hepes 10, Glucose 10, (pH 7.4 with KOH) results in an increase of the coumarin-to-oxonol signal ratio as the oxonol moves away from the coumarin and FRET decreases, i.e. coumarin signal increases while oxonol signal decreases. The final data are expressed as normalized ratio between the two dyes in a column by column display, as shown in the example displayed in Fig. 1a (Control). In presence of 10 μ M gramicidin, the signal is attenuated or nearly abolished

(Gramicidin), in agreement with expectations that the



K⁺-conducting gramicidin channel would collapse the membrane potential generated by the activity of endogenous K⁺ channels. This calibration defines an end point with which to compare the anticipated effect of BoNT HC channel.

Fig. 1

For the HTS-FRET assay of the HC, we have exploited the fact that the endogenous channels of PC12 cells are essentially impermeant to Cs^+ whereas the HC channel is highly conductive for Cs^+ , as shown in independent measurements using patch clamp recordings⁷. Accordingly, the cells are loaded with oxonol in a solution that contains (in mM) NaCl 160, KCl 4.5, CaCl_2 2, MgCl_2 1, Hepes 10, Glucose 10 (pH 7.4 with NaOH), and stimulated in a Hi Cs^+ medium that contains (in mM) CsCl 164.5, CaCl_2 2, MgCl_2 1, Hepes 10, Glucose 10 (pH 7.4 with CsOH). Representative results are shown in Fig. 1b. An increase in fluorescence ratio was determined in presence of the BoNT HC; this is consistent with the fact that, under these conditions, the BoNT HC channel is the major channel which conducts Cs^+ .

The conditions developed thus far require optimization which is a major current effort in the laboratory; however, the initial results validate the HTS-FRET assays on NGF-treated PC12 cells to screen for blockers of BoNT HC.

KEY RESEARCH ACCOMPLISHMENTS

- A key step in the intoxication by BoNT is the translocation of internalized toxin across intracellular membranes to reach its cytosolic targets. A fundamental discovery was the demonstration that the heavy chain acts as both a channel and a transmembrane chaperone for the light chain protease to ensure a translocation competent conformation during transit from acidic endosomes into the cytosol (*Appendix #1*). Thus, the stage is set for pursuing the identification of channel blockers specific for the BoNT heavy chain.
- A high-throughput screen exploiting the VIPR™-Voltage/Ion Probe Reader, is now a proven strategy for the search of compounds that modulate or block the channel activity of BoNT on NGF-treated PC12 cells.

REPORTABLE OUTCOMES

- Publications

Appendix # 1: Koriazova, L. and Montal, M. Translocation of botulinum neurotoxin light chain protease through the heavy chain channel. **Nature Struct. Biol.** [Published online 2 December 2002; doi:10.1038/nsb879] 10:13-18 (2003).

Appendix # 2: Harms, G.S., Orr, G., Montal, M., Thrall, B.D., Colson, S.D., and Lu, H.P. Probing conformational changes of gramicidin ion channels by single-molecule patch-clamp fluorescence microscopy. **Biophys. J.** 85:1826-1838 (2003).

- Invention Disclosure

UCSD #SD2003-007 entitled: Method for antidotal therapy for botulinum toxin (brief).

Title: The botulinum neurotoxin channel as a validated target for intervention aimed to inhibit the translocation of the light chain into the cytosol and therefore to attenuate the neurotoxicity of botulinum neurotoxins.

Inventor: Mauricio Montal

CONCLUSIONS

The key discovery is that the heavy chain of BoNT acts as both a channel and a transmembrane chaperone for the light chain to ensure a translocation competent conformation during its transit from the acidic endosome into the cytosol – its site of action. The light chain is a Zn^{2+} -metalloprotease that cleaves the protein components involved in synaptic vesicle fusion with the neuronal membrane, thereby abrogating synaptic transmission¹. To accomplish this task, the heavy chain operates as a transmembrane protein-conducting channel²: the channel is occluded by the light chain during transit, and open after completion of translocation and release of cargo, acting intriguingly similar to the protein-conducting/translocating channels of the endoplasmic reticulum (ER), mitochondria, and chloroplasts. This finding has outlined a novel way of thinking about BoNT neurotoxicity shifting the focus of attention on its translocation within cells rather than on the protease activity of the light chain, which is known not to be toxic unless it is internalized.

Current progress focused on screens for blockers of the BoNT HC channel as a target for lead discovery are exciting and encouraging. A major outcome of the proposed program would be a blueprint to uncover blockers of the protein-conducting channel of BoNT which may be developed, in due turn, into prophylactic agents to be administered to military personnel prior to deployment or to civilian populations at risk.

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APPENDICES

Appendix #1

Koriazova, L. and Montal, M. Translocation of botulinum neurotoxin light chain protease through the heavy chain channel. **Nature Struct. Biol.** [Published online 2 December 2002; doi:10.1038/nsb879] 10:13-18 (2003).

Appendix #2

Harms, G.S., Orr, G., Montal, M., Thrall, B.D., Colson, S.D., and Lu, H.P. Probing conformational changes of gramicidin ion channels by single-molecule patch-clamp fluorescence microscopy. **Biophys. J.** 85:1826-1838 (2003).

Translocation of botulinum neurotoxin light chain protease through the heavy chain channel

Lilia K. Koriyazova and Mauricio Montal

Published online 2 December 2002; doi:10.1038/nsb879

Clostridial botulinum neurotoxins (BoNTs) abort the process of neurotransmitter release at presynaptic motor nerve terminals, causing muscle paralysis. An enigmatic step in the intoxication process is the mechanism by which the neurotoxin heavy chain (HC) forms the conduit for the translocation of the light chain (LC) protease across the endosomal membrane into the cytosol, its site of action. Here we investigate the mechanism of LC translocation by using the combined detection of channel currents and substrate proteolysis, the two hallmark activities of BoNT. Our data are consistent with the translocation of the LC through the HC channel and show that the LC protease activity is retrieved in the *trans* compartment after translocation. We propose that the BoNT HC–LC complex embedded in the membrane is a transmembrane chaperone, a dynamic structural device that prevents aggregation and achieves translocation of the LC. In this regard, the complex is similar to the protein conducting/translocating channels of the endoplasmic reticulum, mitochondria and chloroplasts.

Clostridial botulinum neurotoxins (BoNTs) act as sequence-specific endoproteases that cleave unique constituents of the SNARE complex — the synaptic vesicle docking/fusion complex^{1–6}. A widely held view is that BoNTs enter cells *via* receptor-mediated endocytosis². Exposure of the BoNT holotoxin to the acidic pH of the endosomes^{7,8} induces a conformational change of the disulfide-linked dichain toxin, allowing the passage across the membrane of the light chain (LC) protease through the putative heavy chain (HC) channel and into the cytosol. Until now, there has been no direct evidence for the translocation of the LC through the HC channel. The crystal structures of BoNT A⁹ and BoNT B¹⁰ confirmed a tri-modular design of the neurotoxin protein: a receptor binding domain, a translocation domain and a catalytic domain. However, these structures were determined at pH 7 and pH 6 for BoNT A and B, respectively, and represent a single conformation of the inactive holotoxin in the soluble, but not the membrane-embedded, form. They also provide no compelling molecular details about the elusive translocation process.

Here, we use the combined detection of channel currents and substrate proteolysis, the two hallmark activities of BoNTs, to determine if HC channel activity and LC translocation are correlated. We demonstrate that channel activity requires acidification of the medium in the aqueous compartment to which BoNT A is added (*cis*) and a reducing environment in the opposite compartment (*trans*). Under these conditions, we detected the LC protease in the *trans* compartment, consistent with its translocation through the HC channel. CD data indicate that only the unfolded conformation of LC A, the LC of BoNT serotype A, correlates with both channel and protease activities of BoNT A. Together, the results imply that the LC partially unfolds at the endosomal pH, passes through the HC channel, refolds at the interface and dissociates from the HC in the reducing cytosolic milieu where it cleaves its substrate SNAP-25. These

findings suggest a novel modality of transmembrane chaperone action based on protein conducting/translocating channels.

LC occludes the HC channel in unreduced BoNT A

To investigate the mechanism of LC translocation, we relied on the channel activity of BoNT A holotoxin and HC A in lipid bilayers and used single channel recordings^{11–13} as a sensitive readout to monitor whether the LC blocks the HC channel as it passes from the *cis* compartment of the membrane to the *trans* one. Evidence that BoNT A forms channels in lipid bilayers^{14,15} and PC12 cells¹⁶, predominantly after exposure to an acidic pH and only after chemical reduction, has led to the hypothesis that the HC forms channels. Our protocol here was to add BoNT A to the *cis* solution, which was held at pH 7.0 to enhance binding to the gangliosides-containing bilayers. After 2 min, the *cis* solution was acidified ($4.5 \leq \text{pH} \leq 5.5$) while the *trans* compartment was held at pH 7.0. Under these conditions, unreduced holotoxin BoNT A does not form channels (Fig. 1a). In contrast, unreduced native (Fig. 1b) and recombinant (Fig. 1c) HC from BoNT A forms channels similar to those recorded with reduced BoNT A (Fig. 1d). Clearly, the HC is a channel irrespective of redox state. These results raised the intriguing question of whether, for the unreduced holotoxin, the HC channel is occluded by the anchored LC.

Disulfide reduction unblocks the HC channel

To test this inference, we examined the effects of the membrane nonpermeable reductant tris-(2-carboxyethyl) phosphine¹⁷ (TCEP) on BoNT A channel activity. The most telling results emerge by examining the contrasting findings generated by the location of TCEP in the *cis* or the *trans* compartments (Fig. 1). Unreduced holotoxin (data not shown), or BoNT A in the absence of TCEP in the acidified *cis* compartment, does not form channels (Fig. 1a). Presence of TCEP in the *trans* compartment,

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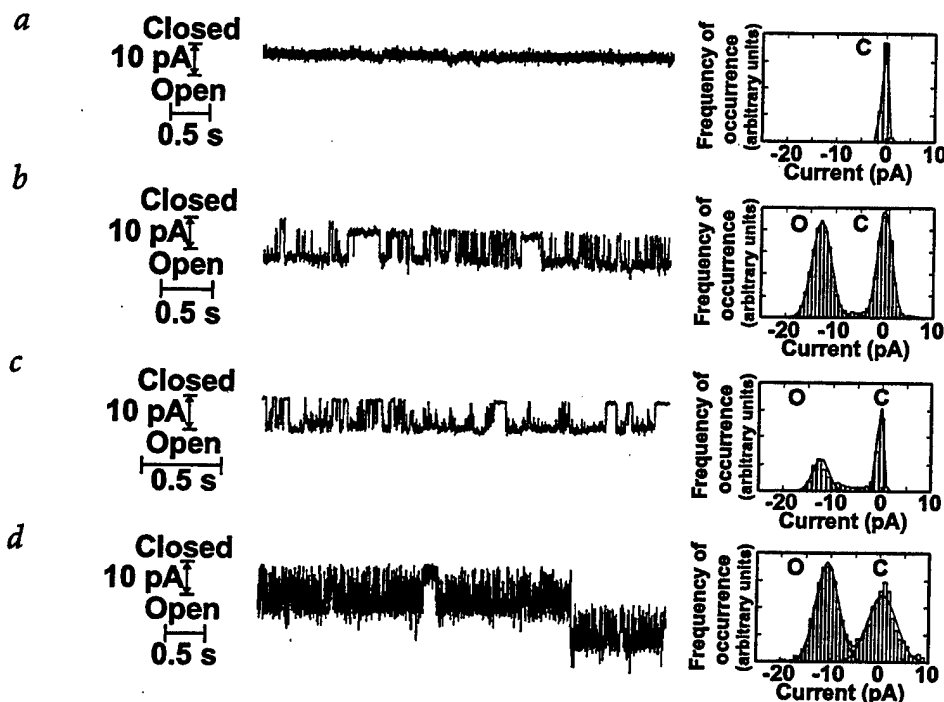


Fig. 1 Channel formation by BoNT A and HCA in lipid bilayers. **a**, Unreduced BoNT A does not form channels ($n = 5$). Records were obtained at -100 mV. The solution in both *cis* and *trans* compartments contained 0.5 M KCl, 1 mM CaCl_2 and 5 mM HEPES, pH 7.1 ± 0.1 . During the experiment, the solution in the *cis* chamber was acidified with 0.8 mM citrate; at the end of the experiment, the final pH was verified to fit within the range $5.0 \leq \text{pH} \leq 5.5$. TCEP (0.5 mM; Sigma) was added to the *cis* compartment 1 min after acidification. A downward deflection indicates channel opening. A current amplitude histogram and Gaussian fit calculated from the record are shown on the right panel; C denotes closed state. Single channel currents from bilayers containing **b**, native HCA or **c**, recombinant HCA. The corresponding current amplitude histogram and Gaussian fit calculated from the record in which a single channel underwent transitions between closed (C) and open (O) states are shown on the right panel. The cumulative probability of the channel being open, P_o , or closed, P_c , is calculated from the area under the corresponding Gaussian curve. For these records, $\gamma = 128$ pS and 96 pS, and $P_o = 0.55$ and 0.48 , respectively (P_o for native and recombinant HC, respectively). The mean $\gamma = 110 \pm 12$ pS and $n = 10$, and $\gamma = 108 \pm 11$ pS and $n = 3$ (P_o for native and recombinant HCA, respectively). Other conditions were as in (a). **d**, Single channel currents of reduced native BoNT A displaying the 110 pS channels. The *trans* compartment contained 0.5 mM TCEP. This segment illustrates the occurrence of two channels. For this record, $P_o = 0.56$, the mean $\gamma = 110 \pm 10$ pS and $n = 10$. A subconductance state with $\gamma \sim 20$ pS is readily identifiable. Protein concentration was $1 \mu\text{g ml}^{-1}$ for (a) and (d), and $5 \mu\text{g ml}^{-1}$ for (b) and (c). Other conditions were as in (a).

however, elicits channel activity (Fig. 1d). When reduced holotoxin is added to the *cis* compartment, followed by acidification, channels are formed and there is no need for TCEP in the *trans* compartment. Notably, addition of TCEP to the *cis* compartment 1 min after acidification fails to evoke channel activity (Fig. 1a). The absence of channel activity when TCEP is added to the *cis* compartment after acidification is interpreted as indicative of the translocation of the LC through the HC channel, rendering the critical disulfide link between LC and HC inaccessible to the membrane-impermeant TCEP. Presumably, the LC remains attached to the HC, occluding the HC channel, whereas reduction in the *trans* compartment releases the LC, thereby unblocking the HC channel. This observation implies that the LC blocks the HC channel during its translocation.

The LC selectively blocks the HC channel

This prediction was tested using isolated HC and LC and addressing the specific effect of the LC on HC channels depending on its location in the *cis* or the *trans* compartments (Fig. 2). Neither native nor recombinant LC A forms channels^{14,15}. A ten-fold molar excess of LC in the acidified *cis* compartment, in the absence of reductant, does not affect the HC channel activity (Fig. 2a). In contrast, a stoichiometric amount of LC in the neutral *trans* compartment attenuates and eventually silences the

HC channel activity (Fig. 2b): the single channel conductance (γ) is sharply reduced (Fig. 2b inset) from 110 pS to 25 pS, and the channel openings become brief, appearing as transient spikes at this time resolution. This pattern is characteristic of open channel blockers and arises from the progressive shortening of the residence time in the open state¹². Blocking is elicited by both native and recombinant LC on both the native and recombinant HC channels, is independent of the polarity of the applied voltage and is not produced by thermally or guanidine-HCl-denatured and inactivated LC (Fig. 2c). Thus, at pH 7.0 , the LC selectively blocks the HC channel.

LC translocation revealed by its protease activity

To examine if the LC protease goes through the HC channel, we developed a high-sensitivity ELISA and scaled up the single channel measurements to detect numerous ($\geq 1,000$) channels (Fig. 3a). The *trans* compartment contained the LC substrate, SNAP-25 (refs. 2,3). A period of 15 – 30 min was fixed for each assay; afterward, the entire volume of the *trans* chamber was collected and assayed for SNAP-25 proteolysis. No cleavage of SNAP-25 was detected in the absence of BoNT A (control) or in the presence of LC A in the *cis* compartment (Fig. 3c). The latter control indicated that the bilayer prevented mixing of isolated LC (*cis*) with SNAP-25 (*trans*) and confirmed that LC *per se* can-

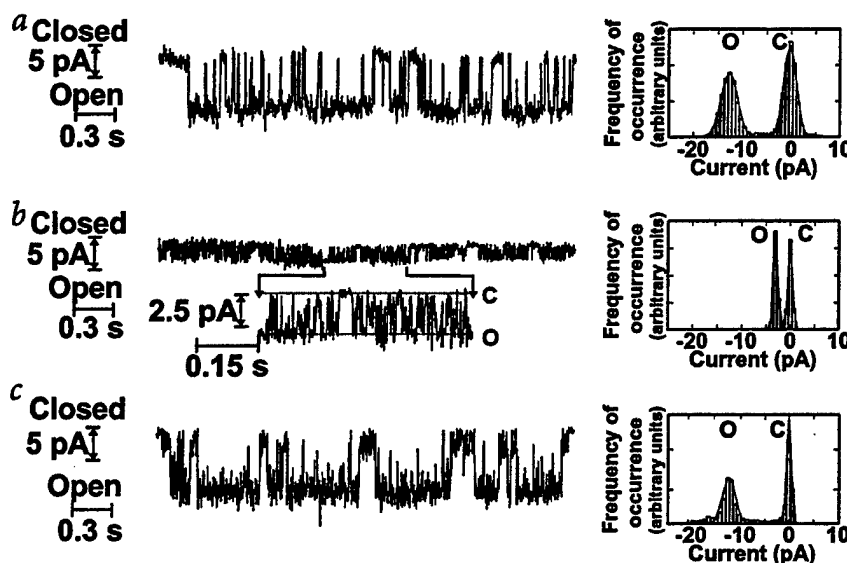


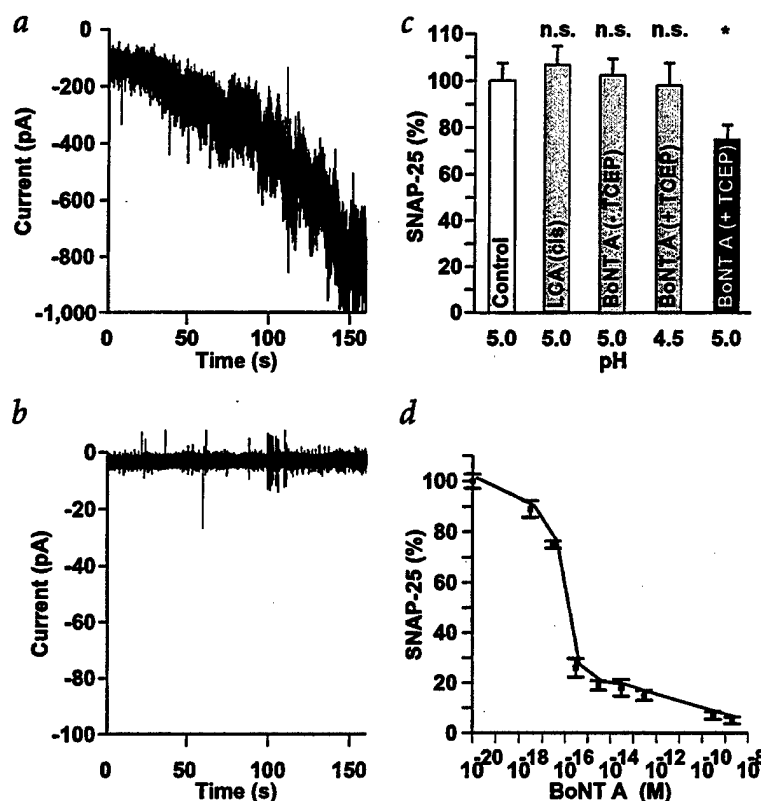
Fig. 2 Blocking of HC A channels by the LC A. **a**, Single channel currents of native BoNT HC in the presence of LC in the *cis* chamber. LC A ($25 \mu\text{g ml}^{-1}$) was added after the onset of HC A channel activity. For this record, $\gamma = 124 \text{ pS}$ and $P_o = 0.56$. The mean $\gamma = 113 \pm 9 \text{ pS}$ and $n = 5$. **b**, Single channel currents of native BoNT HC in the presence of LC in the *trans* chamber. LC A ($2.5 \mu\text{g ml}^{-1}$) was present from the beginning of the experiment. The segment delimited by the arrows is displayed at higher resolution. For this record, $\gamma = 29 \text{ pS}$ and $P_o = 0.51$. The mean $\gamma = 25 \pm 4 \text{ pS}$ and $n = 5$. Note the occurrence of fast transitions between open and blocked states. **c**, Single channel currents of native BoNT HC in the presence of denatured LC A ($10 \mu\text{g ml}^{-1}$) in the *trans* chamber. LC A was inactivated by incubation in 6 M guanidinium-Cl for 30 min before the addition. Inactivation was confirmed by direct assay of SNAP-25 cleavage. For this record, $\gamma = 123 \text{ pS}$ and $P_o = 0.54$. The mean $\gamma = 112 \pm 9 \text{ pS}$ and $n = 5$. Protein concentration was $5 \mu\text{g ml}^{-1}$ for HC. Other conditions were as in Fig. 1a.

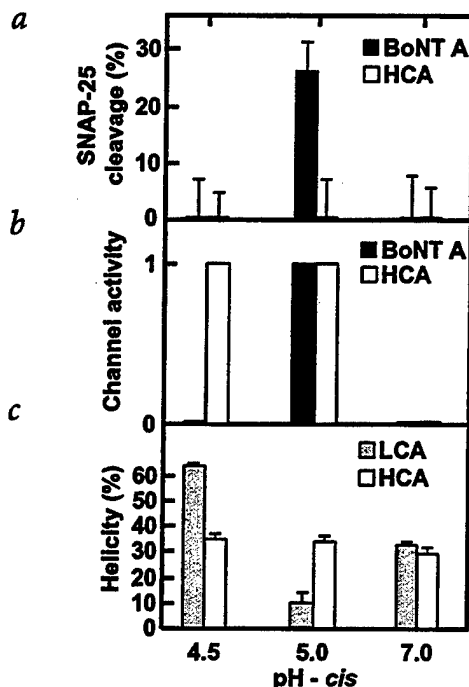
not permeate through an unmodified lipid barrier. Cleavage of SNAP-25 requires the presence of TCEP on the *trans* side and pH 5.0 on the *cis* side (Fig. 3c), a condition that correlates with the insertion of multiple channels in the bilayer (Fig. 3a, equivalent to Fig. 1a for single channels). This correspondence argues that there is proteolytic activity of the LC on the *trans* side only under conditions in which the channel activity of the holotoxin is detected. This is consistent with the concomitant absence of channel (Fig. 3b) and LC protease (Fig. 3c) activity when the pH on the *cis* side was 4.5.

To estimate the number of LC molecules translocated across the bilayer, we

established a calibration curve (Fig. 3d). The minimum BoNT A LC activity was discerned at $3 \times 10^{-18} \text{ M}$, producing a $10 \pm 5\%$ cleavage of SNAP-25. At $3 \times 10^{-17} \text{ M}$, $25 \pm 5\%$ cleavage was reliably measured. This value approximates the value detected with TCEP on the *trans* side and at pH 5.0 on the *cis* side (Fig. 3c),

Fig. 3 Detection of the endopeptidase activity of translocated LC A in bilayer experiments. **a**, Time course of insertion of multiple BoNT A channels into a membrane at pH 5. Records obtained at -10 mV . The *cis* solution was 0.1 M KCl , 1 mM CaCl_2 and 5 mM HEPES , pH 7.1 ± 0.1 . After addition of BoNT A ($0.25 \mu\text{g ml}^{-1}$) the pH was adjusted to 5.0. The *trans* solution was 0.01 M KCl , 1 mM CaCl_2 , 5 mM HEPES , pH 7.1 ± 0.1 , $4 \mu\text{M}$ N-terminal biotinylated SNAP-25, 0.5 mM TCEP and $10 \mu\text{M Zn(CH}_3\text{COO)}_2$. **b**, No detectable channel activity of BoNT A at pH 4.5. Conditions identical to (a) except that the pH in the *cis* compartment was 4.5. **c**, SNAP-25 cleavage in samples collected from the *trans* chamber of bilayer experiments equivalent to the representative records shown in (a) and (b). Error bars indicate s.d. and $n = 6$ experiments, except for BoNT A (+TCEP) at pH 5.0 (a), for which $n = 10$. Probabilities are indicated by the asterisk from t-tests performed with respect to control for each experimental condition separately; 'n.s.' denotes not significant; asterisk, $P < 0.001$. **d**, SNAP-25 cleavage as function of BoNT A concentration determined by ELISA ($n = 4$). The values obtained for the isolated LC A at selected concentrations agree well with those shown in (d).





which corresponds to ~3,600 LC molecules in the experimental samples. This number is in close agreement with the maximum number of channels inserted in the bilayer under these assay conditions, ~2,000 (Fig. 3a), as estimated from the macroscopic membrane conductance $\sim 1 \times 10^5$ pS and $\gamma = 50$ pS measured in 0.1 M KCl. Together, these findings demonstrate translocation of the LC through the HC channel and retrieval of the LC protease activity at neutral pH.

How does the LC go through the HC channel?

The diameter of the BoNT A or HCA channel, estimated from γ , is ~15 Å (refs. 14,18), raising the question of how a pore of this size could accommodate the globular LC with dimensions of $55 \text{ Å} \times 55 \text{ Å} \times 62 \text{ Å}$ (ref. 9). These considerations imply that the LC unravels at the endosomal pH ($5.1 \leq \text{pH} \leq 5.4$) to go through the channel and refolds after exiting at the neutral cytosolic pH. To assess the pH-induced structural changes of the LC and HC associated with their respective protease (Fig. 4a) and channel (Fig. 4b) activities, we used CD spectroscopy (Fig. 4c). At pH 7.0, the α -helical content of the individual LCA and HCA is comparable to that calculated from the crystal structure of BoNT A⁹. The α -helical content of the HCA was unaffected by pH. For LCA, there is a sharp decrease in helicity at pH 5.0 (Fig. 4c), which correlates with the occurrence of channel (Figs. 1a, 3a, 4b) and protease (Figs. 3c, 4a) activities of BoNT A. In contrast, at pH 4.5, there is a drastic increase in LC helicity coincident with the absence of channel (Figs. 3b, 4b) and protease (Figs. 3c, 4a) activities. Together, the data indicate that only the unfolded conformation of LCA correlates with both channel and protease activities of BoNT A. This suggests that, under the conditions prevalent in the cell after BoNT A entry — specifically, the endosome/cytosol pH gradient⁷ and a cytosolic reducing milieu¹⁹ — the LC unfolds or partially unfolds in the endosome, goes through the HC channel, dissociates and refolds at the interface, and cleaves SNAP-25 in the cytosol. At pH 4.5, the acid-induced conformational change of BoNT A allows insertion of the HC

Fig. 4 BoNT A endopeptidase activity correlates with BoNT A channel activity and the unfolding of LC A. **a**, Endopeptidase activity in samples collected from the *trans* chamber of bilayer experiments (Fig. 3c) as function of pH in the *cis* compartment. The *trans* compartment pH was 7.0. **b**, Channel activity of BoNT A and HCA as function of pH in the *cis* compartment; the *trans* compartment pH was 7.0. Absence or presence of channel activity is arbitrarily defined as 0 or 1; $n = 6$ for HCA and $n = 10$ for BoNT A (Fig. 3a,c). Other conditions equivalent to those of Fig. 3a. **c**, α -helical content of LCA and HCA as function of pH calculated from far UV-CD measurements carried out at 25 °C ($n = 3$).

into the membrane, but the high helical content of the LC traps it within the HC channel and occludes the channel, thereby precluding detection of channel or protease activities.

The HC acts as a protein-translocating channel

The interplay between an unfolded LC embedded within the HC is reminiscent of the maintenance of an unfolded or partially folded state of polypeptides by chaperones. Is the HC acting as a transmembrane chaperone for the LC to ensure a translocation-competent conformation, with its channel activity simply the expression of an empty state? Three protein analogies are noteworthy: prefoldin²⁰, protein disulfide isomerase (PDI)²¹ and *Salmonella* SicP²². Neither prefoldin nor PDI require ATP for their chaperone activity. The prefoldin hexamer consists of six long α -helical coiled coils assembled to a double β -barrel template. The distal segments of the coiled coils generate the hydrophobic surface required for binding of unfolded polypeptides. The crystal structure of prefoldin²⁰ shows similarity to the long helices of the translocation domain of BoNT A⁹ and B¹⁰ (Z score = 5.0, r.m.s. deviation of 4.5 Å and 3.7 Å, respectively²³). PDI, whose substrate-binding cycle is regulated by its redox state, is crucial to the unfolding of cholera toxin and for its translocation from the ER lumen into the cytosol of target cells²¹. Numerous bacterial pathogens deliver proteins (effectors) into target cells with the assistance of specific chaperones. To fit through the secretion 'channel', with an apparent cut-off diameter of 30 Å, the effector is kept unfolded by interacting with the chaperone. The 1.9 Å crystal structure of the chaperone-binding domain of the *Salmonella* effector protein SptP in complex with its cognate chaperone SicP shows the unfolded conformation of SptP threaded around three chaperone molecules²². For BoNT A, we may envision a chaperone activity driven by a pH gradient across the endosome in which the HC channel prevents the aggregation of the LC in the acidic vesicle interior *via* hydrophobic interactions, thereby maintaining the unfolded conformation during translocation and releasing the LC after it refolds at the neutral cytosolic pH. In this process, the HC channel is occluded by the LC during transit and opens after completion of translocation and release of cargo. In other words, the BoNT A HC-LC complex embedded in the membrane is a dynamic structural device that prevents aggregation and achieves translocation of the LC. An analogous mechanism has been invoked for the translocation of the catalytic domain of diphtheria toxin²⁴. There is a close resemblance of the occluded BoNT A channel with the translating ribosome-protein-conducting channel complex (yeast ER sec61 complex): the cryo-EM reconstruction at 15.4 Å resolution led to the view that the nascent polypeptide chain is tightly accommodated within the channel, hindering conductance²⁵. Given a constriction of ~15 Å, the channel may allow passage of only α -helices²⁵. The 8 Å structure of the sec61-related bacterial protein translocation complex SecYEG reveals a deep cavity at the dimer interface that is occluded at the periplasmic side and may represent the closed translocation channel²⁶. Similarly, unfolding of precursor proteins by the

mitochondrial import system is required for translocation across the mitochondrial Tim23 channel, which has a pore diameter of ~13 Å (ref. 27). For chloroplasts, the peptide-translocating channels of the outer (Toc75)²⁸ and inner (Tic110)²⁹ membranes display pore diameters of ~14 Å and 15 Å, respectively. Protein conducting/translocating channels seem, therefore, to constitute a novel mode of chaperone action. Our findings also suggest that the BoNT channel may represent a potential target for intervention to attenuate BoNT neurotoxicity.

Methods

Reconstitution of BoNT channels in lipid bilayers. Lipid bilayers were formed at 25 ± 1 °C by apposition of two monolayers³⁰, either at the tip of patch pipets¹³ for single channel recordings (Figs. 1, 2) or across a 500 µm aperture on a teflon partition between two aqueous compartments (250 µl) for LC translocation experiments¹² (Figs. 3, 4). The lipid composition was optimized to enhance the binding and insertion of BoNT A and HC and consisted of (1,2-diphytanoyl-sn-glycero-3)-phosphatidylethanolamine, -phosphatidylcholine and -phosphatidylserine (Avanti Polar Lipids), supplemented with GT1b gangliosides (Fidia Research Laboratories) at a weight ratio 2:2:1:0.125 for the four lipids used. Acquisition and analysis of channel currents were essentially as described^{12,13}. Throughout the text, data are shown as mean ± s.d. and *n* = number of experiments.

Native BoNT A holotoxin, LC and HC. Purified native BoNT A, LC A and HC A were from M.C. Goodnough (University of Wisconsin). BoNT A nicking was estimated by SDS-PAGE as ≥99%.

Recombinant BoNT A LC and HC. BoNT A LC clone in pBN3 expression vector was obtained from H. Niemann. The LC protein was expressed in the *Escherichia coli* strain BL21 (Novagen) and purified essentially as described³¹. The recombinant HC clone was obtained from B.R. Singh (University of Massachusetts) and subcloned into pCRT7/CT TOPO TA vector (Invitrogen). The HC protein was expressed in the *E. coli* strain BL21-CodonPlus(DE3)-RIL (Stratagene) and purified from the 6 M guanidinium-Cl–clarified inclusion bodies extract by nickel chelate chromatography in presence of 0.05% dodecylphosphocholine (Avanti Polar Lipids).

Endopeptidase activity of translocated LC in bilayer experiments. Recombinant SNAP-25 was expressed and purified as described³². SNAP-25 truncation was evaluated by ELISA. N-terminal

biotinylation of SNAP-25 was accomplished at pH 7.2 using the *N*-hydroxysuccinimide ether of biotin (biotin-XX-NHS, Calbiochem). Cleavage of biotinylated SNAP-25 (4 µM) proceeded for 24 h at 37 °C. ELISA microtitration plates (Dynex Technologies) were coated with goat polyclonal antibodies against SNAP-25 C-terminal peptide (Santa Cruz Biotechnology). Plates were blocked with biotin-free 1% casein colloid buffer (Research Diagnostics). Each bilayer experimental sample was tested in duplicate and in parallel to control wells on the same plate, after incubation for 1 h at 37 °C. Streptavidin-polymerized horseradish peroxidase (Poly-HRP-SA₉₀) conjugate (Research Diagnostics) was applied at 1:5,000 dilution. Absorbance at 405 nm was determined after addition of ABTS (2,2'-azino-bis-3-ethylbenzothiazoline-6-sulfonic acid; Boehringer Mannheim), using an automatic microplate reader (EL311; Bio-Tek Instruments). On the basis of the standard curve (*n* = 4), the ELISA detection limit of <3 × 10⁻¹⁸ M was established as the BoNT concentration corresponding to 3 s.d. from the maximum (100%) uncleaved SNAP-25 (Fig. 3d)³³.

CD spectroscopy. CD measurements were taken at 25 °C on an AVIV 202 CD spectrometer (Aviv Instruments) using a 0.1 cm path length cell and analyzed as described³⁴. For all spectra, the average of three scans was obtained. To correlate the CD results with the bilayers measurements, we used 100 mM KCl, 50 mM potassium phosphate buffered to pH 7.0, 5.0 or 4.5. These buffers were supplemented with 1 mM β-mercaptoethanol and 0.05% dodecylphosphocholine to prevent aggregation, which increases with acidification. The differences between our CD results and other reports³⁵ likely arise from the different experimental conditions used. Li and Singh³⁵ used 50 mM NaCl buffered with 50 mM Tris for the pH 7.0 measurements and 50 mM NaCl buffered with 20 mM Tris for the pH 4.7 measurements, and their measurements were conducted at 4 °C.

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Competing Interests statement

The authors declare that they have no competing financial interests.

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Probing Conformational Changes of Gramicidin Ion Channels by Single-Molecule Patch-Clamp Fluorescence Microscopy

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ABSTRACT Complex conformational changes influence and regulate the dynamics of ion channels. Such conformational changes are stochastic and often inhomogeneous, which makes it extremely difficult, if not impossible, to characterize them by ensemble-averaged experiments or by single-channel recordings of the electric current that report the open-closed events but do not specifically probe the associated conformational changes. Here, we report our studies on ion channel conformational changes using a new approach, patch-clamp fluorescence microscopy, which simultaneously combines single-molecule fluorescence spectroscopy and single-channel current recordings to probe the open-closed transitions and the conformational dynamics of individual ion channels. We demonstrate patch-clamp fluorescence microscopy by measuring gramicidin ion channel conformational changes in a lipid bilayer formed at a patch-clamp micropipette tip under a buffer solution. By measuring single-pair fluorescence resonance energy transfer and fluorescence self-quenching from dye-labeled gramicidin channels, we observed that the efficiency of single-pair fluorescence resonance energy transfer and self-quenching is widely distributed, which reflects a broad distribution of conformations. Our results strongly suggest a hitherto undetectable correlation between the multiple conformational states of the gramicidin channel and its closed and open states in a lipid bilayer.

Subtle conformational changes of ion channels play an important role in regulating the function and dynamics of ion channels (Karlin, 2002; Madden, 2002; Choe, 2002). Channel conductance recording analysis and kinetic modeling (Cross et al., 1999; Andersen et al., 1999), together with site-directed mutagenesis, protein modifications, and static structural analysis provide extensive insights into the conformations and the conformational rearrangements associated with channel activity (Karlin, 2002; Madden, 2002; Choe, 2002; Glauner et al., 1999; Cha et al., 1999). However, the conformational dynamics underlying the mechanisms of ion channel action are still largely unknown due to lack of direct measurements. Mechanistic characterization of the conformational changes has been primarily based on kinetic model analyses of the electric current trajectories obtained by single-channel and whole-cell patch-clamp recordings that are not sensitive to subtle conformational intermediate states, including electrically undetectable “silent” conformational states. Protein dynamics may involve significant static and dynamic inhomogeneities (Lu et al., 1998; Xie and Trautman, 1998). For instance, ion channel dynamics can be modulated by inhomogeneous local membranes and a fluctuating environment. These static and dynamic inhomogeneous conformational dynamics are extremely difficult to study by conventional experimental approaches.

Recently, our group (Orr et al., 2001, 2002; Harms et al., 2002, 2003) and others have made significant progress in developing new approaches to studying ion channels using both fluorescence imaging and electric patch recording. Utilizing a micro-pinhole patch technique, Yanagida and coworkers (Ide and Yanagida, 1999; Ide et al., 2002) first reported fluorescence images acquired before or after a single-channel current measurement on the same sample lipid bilayer. Woolley, Schuetz, and their co-workers (Loughheed et al., 2001; Borisenko et al., 2003) have elegantly demonstrated the feasibility of combining single-molecule fluorescence imaging and electric recording on gramicidin ion channels. Isacoff and co-workers (Sonnleitner et al., 2002) have reported a single-molecule imaging and whole-cell patch recording study of structural rearrangement of voltage-gated Shaker K⁺ channels in the plasma membrane of living cells. A related experimental effort by Schmidt and his co-workers (Harms et al., 2001) has demonstrated wide-field fluorescence microscopy imaging of the diffusion and aggregation of yellow-fluorescent protein-labeled L-type Ca²⁺ channels in living cells. Another related experiment by Zagotta and his co-workers (Zheng and Zagotta, 2000) has studied gating rearrangements of cyclic nucleotide-gated channel proteins by combined fluorescence imaging and patch-clamp recording at multiple-channel-averaged measurements. These research efforts have begun to provide a new paradigm for studying ion channel conformational dynamics and mechanisms and for obtaining information not obtainable by the conventional patch current recording measurements, a primary experimental approach for decades.

Here, we demonstrate a new approach, patch-clamp fluorescence microscopy (PCFM), by combining single-molecule fluorescence imaging of fluorescence resonant energy transfer (FRET) and fluorescence self-quenching

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with a single-channel electric current patch recording. Using PCFM, we were able to correlate in real-time the single-channel open-closed kinetics with simultaneous changes in optical signals associated with conformational changes in a gramicidin ion channel in a lipid bilayer. By utilizing a typical patch-clamp technique in this new approach, the application of PCFM for studying ion channel dynamics in living cells (Orr et al., 2002) is now feasible. A two-state kinetic model has been widely referenced to illustrate the gramicidin ion channel kinetic behavior, i.e., the association and dissociation of the gramicidin dimer corresponding to the channel's open and closed states. However, the nature of the conformational changes and the dynamics that regulate channel activity are still largely unclear, since there are no methods of providing a direct real-time single-molecule measurement of both ion channel electric current and conformational changes. Our new experimental results, not obtainable by using ensemble-averaged methods or single-channel patch electric recording alone, suggest the occurrence of multiple intermediate conformational states underlying the gramicidin channel dynamics.

MATERIALS AND METHODS

Linear gramicidin produced naturally from the bacterium *Bacillus brevis* (Hladky and Haydon, 1970) is a typical "model" ion channel (Koepppe and Andersen, 1996; Cross et al., 1999; Andersen et al., 1999), and its conductive open state is presumably a dimer (Veatch et al., 1975; Koepppe and Andersen, 1996; Cross et al., 1999; Andersen et al., 1999) of two 15-amino acid peptides forming β -helices that meet head-to-head at the N-termini in the interior of a lipid bilayer (Koepppe and Andersen, 1996; Cross et al., 1999; Andersen et al., 1999; Roux and Karplus, 1994). Because of the available extensive knowledge on the gramicidin ion channels in lipid bilayers, we chose this system to demonstrate our combined single-molecule approach of PCFM.

Fig. 1 shows the experimental setup for the PCFM single-channel electric current recording and fluorescence imaging. We recorded the single-channel current of dye-labeled gramicidin dimers that were incorporated in lipid bilayers formed at the tip of a patch pipette (Montal and Mueller, 1972; see Fig. 1, *inset*) as we simultaneously imaged the pipette tip using wide-field single-molecule fluorescence microscopy (Schmidt et al., 1996; Ma et al., 2000; Han et al., 2001; Bartko et al., 2002; Seisenberger et al., 2001; Deschenes and Vanden Bout, 2001; Cognet et al., 2000).

Single-molecule fluorescence microscope

A microscope (Nikon Diaphot 300, Nikon, Kanagawa, Japan) equipped with a 60 \times objective (N.A. = 1.4) and 4 \times relay lens was used in this work. In all experiments, the illumination intensity was set to 5 kW/cm² for tetramethylrhodamine (TMR)-gramicidin at 514 nm (Ar-ion laser, Molecular Probes, Eugene, OR; MWK Laser, Riverside, CA) or 532 nm (Verdi 10 W diode laser, Coherent, Palo Alto, CA), and 1.5 kW/cm² for Cy5-gramicidin at 632 nm (HeNe laser, Uniphase or dye laser, CR-5999, Coherent). The polarization of the excitation light was controlled by using a $\lambda/4$ waveplate (Meadowlark Optics, Frederick, CO). Filter combinations for TMR consisted of DCLP555, HQ625/100 (Chroma Technology Corp., Brattleboro, VT) and OG550 (Schott, Duryea, PA), and for Cy-5 of DCLP645, HQ690/90, and RG645 (Schott). The single-pair fluorescence resonant energy transfer (spFRET) measurements were done using a custom-built dual-dichroic beam-splitter (a combination of a DCLP555 and DCLP645, Chroma Technology Corp.) and band-pass filter in addition to either OG550 or RG645, permitting the detection of single fluorophores by a nitrogen-cooled CCD camera (Spec-10 1340 \times 400B or 700B, Roper Scientific, Trenton, NJ). The microscope system total detection efficiency was $10 \pm 2\%$ for TMR and $8 \pm 2\%$ for Cy5. We estimated the total detection efficiency from the collection efficiency of the objective, the spectral throughput of the filter set, and the spectral detection efficiency of the CCD camera. We also subtracted the estimated reflections for each glass surface in the optical beam path. This device ensured the toggling of the excitation wavelength between 532 nm and 632 nm, and the detection of the emission at $8 \pm 2\%$ efficiency for TMR and $5 \pm 2\%$ efficiency for Cy5 with the FRET filters. The images were taken using 5-ms exposures at 1–10 Hz.

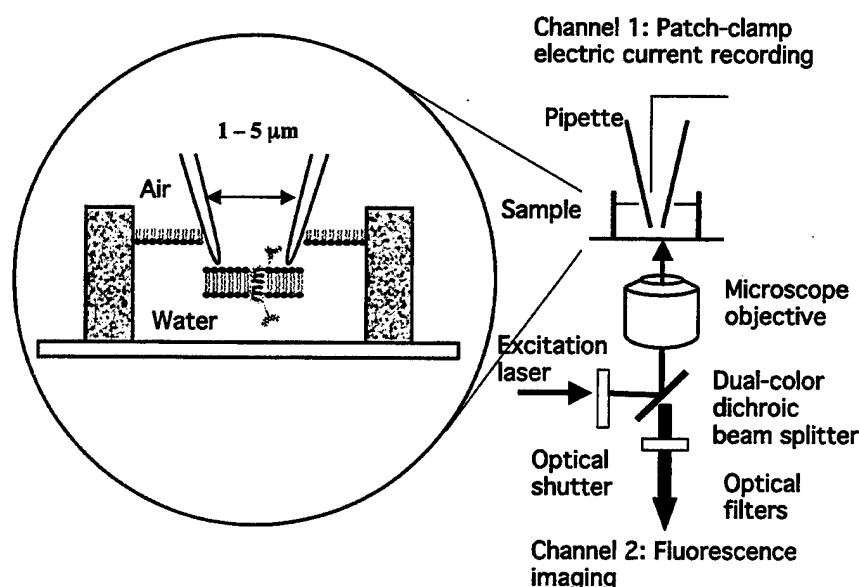


FIGURE 1 Experimental apparatus and measurement configuration of PCFM. Lipid bilayers were formed at the tip of a patch pipette by apposition of two monolayers of mixed lipids (4:1 dPhPE:DPhPC). Dye-labeled gramicidin monomers are introduced from both sides of the bilayer, and single-channel electric currents are recorded by using typical patch-clamp instrumentation. The laser excitation was focused through a microscope objective on the pipette tip, and the fluorescence was imaged on a CCD camera.

Lipid bilayer formation at the tip of a patch-pipette

Lipid bilayers were formed at the tip of patch pipettes by the apposition of two monolayers (Montal and Mueller, 1972), using 4:1 diphytanyolphosphatidylethanolamine (dPhPE):diphytanyolphosphatidylcholine (dPhPC) (Avanti, Alabaster, AL). Gramicidin dissolved in ethanol (10^{-9} M) was added to the aqueous subphase (1 M KCl, 1 mM CaCl_2 , and 5 mM Hepes of pH 7.5) after bilayer formation (Fig. 1, *inset*). Gramicidin monomers were labeled with either TMR or Cy5. Homodimers of TMR-labeled gramicidin were assembled by adding gramicidin-TMR monomers to both sides of the membrane; gramicidin-TMR-gramicidin-Cy5 heterodimers were generated by introducing TMR-gramicidin and Cy5-gramicidin from opposite sides of the bilayer (O'Connell et al., 1990). The latency period to form an active channel is typically a few minutes after gramicidin insertion into the bilayer and presumably arises from the slow diffusion of individual gramicidin monomers to form a dimer.

Single-channel electric current patch recording

Single-channel electric currents elicited at constant applied voltage (-100 – $+100$ mV) were recorded using an EPC-9 amplifier (HEKA Electronics, Lambrecht, Germany). The recorded channel electric signal was sampled at 4 kHz and filtered at 3 kHz. Control experiments using unlabeled gramicidin A revealed distributions of current amplitude and open/closed dwell-time similar to those of the dye-labeled gramicidin. The dye-labeling of the gramicidin C-terminal did not significantly alter the gramicidin ion channel activities (Lougheed et al., 2001; Borisenko et al., 2003). The presence of a single channel at the patch was confirmed by recorded channel-current trajectories showing typical single-channel open-closed behavior (O'Connell et al., 1990; Sawyer et al., 1989; Veatch et al., 1975). The following criteria were applied to ensure that the recorded channel activity was from a single ion channel and not due to contamination or membrane instability: 1), no channel activity was detected before adding gramicidin; 2), only bursts of channel activity were considered; 3), a high membrane resistance (at the giga-ohm range) leading to a high signal-to-noise ratio (S/N) was achieved; and 4), membrane breakdown occurred when 300–500 mV polarization voltage was applied to the patch. These criteria are also practiced commonly in other labs (O'Connell et al., 1990; Sawyer et al., 1989; Veatch et al., 1975; Sigworth et al., 1987) to ensure a single gramicidin channel patch recording. About 10% of all the attempts were successful in incorporating a single gramicidin channel at the patch; membranes without channel activity or with more than one channel were not pursued.

Dye labeling of gramicidin monomers

Gramicidin C with Gly and Lys residues attached to the C-terminal (synthesized by Genemed Synthesis, Inc.) was used in this work. TMR or Cy5 HNS esters were covalently bonded by acylation to the ϵ -amino group of the Lys residue. The full sequence of the gramicidin used for the fluorescence labeling was Formyl - V - G - A - (d)L - A - (d)V - V - (d)V - W - (d)L - Y - (d)L - W - (d)L - W - G - K-NHCH₂OH. Multiple HPLC purifications (C8 column with 80% methanol solvent) ensured the purity of the dye-labeled gramicidin samples, which was further confirmed by MALDI-TOF mass spectroscopy (molecular weight of gramicidin C-TMR, calculated = 2412 and measured = 2415; molecular weight of gramicidin C-Cy5, calculated = 2637 and measured = 2641).

Fluorescence imaging data analyses

The fluorescence background was subtracted by averaging photon counts for each pixel or by fitting the background to a wide 2-D Gaussian. The

individual fluorescent "hot spots" were identified by the maxima of a 2-D Gaussian-correlation filter (Schmidt et al., 1996). The region around these maxima was fitted to a 2-D Gaussian, reporting the intensity, width, position, and errors of the least square fit (Schmidt et al., 1995, 1996; Press et al., 1990).

Single-molecule fluorescence control experiments

Single-molecule imaging was conducted of dye-labeled gramicidin in Langmuir-Blodgett (LB) bilayers and spin-coated over polymethyl methacrylate (PMMA) polymer-coated cover slips. The single-molecule fluorescence imaging signal was confirmed by several typical verification measurements, including observations of single-step photobleaching events (Fig. 2 *a*, *inset*) and of linearly polarized dipoles (data not shown). The dPhPE:dPhPC 4:1 was doped with 10^{-8} mol TMR-gramicidin C or Cy5-gramicidin C per mol lipid and were deposited at 22°C and 28 dynes/cm on glass slides using LB deposition. Single-molecule fluorescence imaging measurements, using essentially the same experimental parameters and configurations that were used to image the bilayers containing individual dye-labeled gramicidin molecules at the tip, provided a quantitative

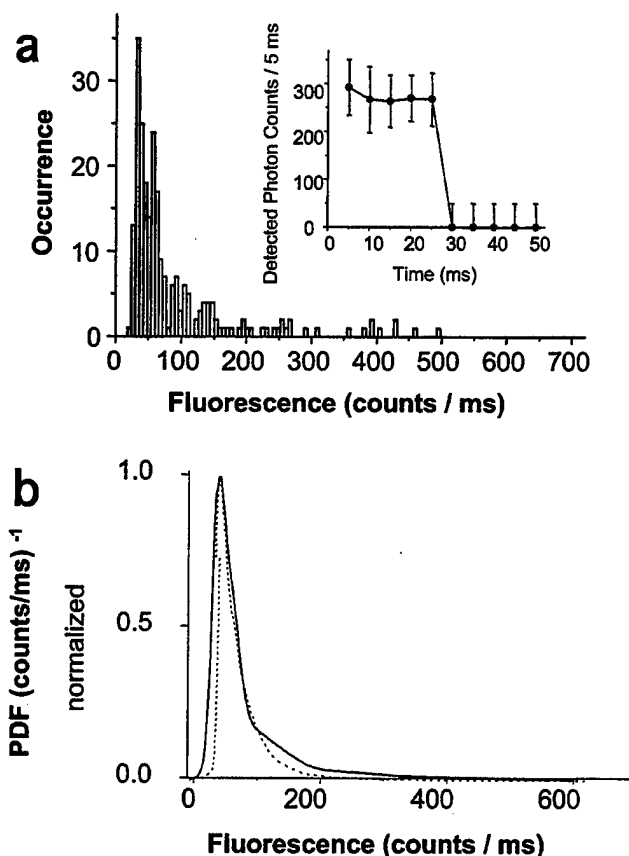


FIGURE 2 (*a*) Histogram of single hot-spot fluorescence intensities deduced from 2-D Gaussian fitting to each spot imaged at the patch-clamp tip. Inset: intensity trajectory from a "hot spot" image of a single TMR-gramicidin molecule. (*b*) The probability density function (*normalized*) deduced from the histogram in *a*, and from the histogram of hot-spot intensities on glass-supported lipid bilayers (*dotted line*) as a control. Our control experiments and those of others (Lougheed et al., 2001) show no observable perturbation by dye labeling on the gramicidin channel activity.

reference for the single-molecule photon count rates under the wide-field imaging configuration (Fig. 2 *a*) in our experiments, which is a typical calibration commonly practiced in single-molecule imaging experiments (Schmidt et al., 1995, 1996; Seisenberger et al., 2001; Harms et al., 2001). The single-molecule images were analyzed by the least-square 2-D Gaussian fitting, as discussed above.

The probability density function (PDF) of the photon counts,

$$PDF(x) = \sum_i \frac{w_i}{\Delta I_i \sqrt{2\pi}} \exp \left\{ -\frac{(x - I_i)^2}{2\Delta I_i^2} \right\},$$

is constructed using parameters from the single-molecule image photon counts, where I_i , ΔI_i , and w_i represent the photon counts, its standard deviation, and its relative weight. The PDF statistically evaluates the imaging photon counts with their standard deviations and occurrences and provides a more reliable distribution than an occurrence histogram does (Schmidt et al., 1995). Under our set-up and experimental conditions, the average photon count detection rate for the TMR-gramicidin C monomers was 68 ± 5 counts/ms with a PDF peak of 45 ± 5 counts/ms and PDF width of 32 counts/ms. For Cy5-gramicidin C, the average photon count was 46 ± 5 counts/ms with a PDF peak of 36 ± 5 counts/ms and width of 30 counts/ms. The width of the PDF distributions indicates that the level of signal fluctuations is due primarily to shot noise and to the fluorescence quantum yield distribution of the individual dye molecules in a spatially inhomogeneous local environment (Lu and Xie, 1997; Xie and Trautman, 1998).

There are inevitable differences in the fluorescence photon detection efficiency and the signal-to-noise ratio of imaging single molecules on the LB layer or the PMMA polymer, as in a control experiment, and in imaging single molecules at a patch-clamp pipette tip, as in a real PCFM measurement (Fig. 2 *b*). There are at least three reasons for these differences: 1), There are different configurations for the pipette tip under a buffer solution in the PCFM measurement and the cover slip with individual dye-labeled gramicidin molecules in LB layers under water or spin-coated individual dye-labeled gramicidin molecules on PMMA polymer films. These intrinsic differences in the experimental configuration apparently cause different signals and noise levels associated with a change in the refractive index and light-scattering conditions. 2), There are different fluorescence quantum efficiencies under different conditions for the LB bilayer, a polymer, and the lipid bilayer at a pipette tip under a buffer. 3), There are different focal planes of the wide-field laser excitation and fluorescence collection, which are at the upper surface of the cover slip for the control experiment but at ~ 70 μm above the cover slip in the buffer for imaging at the pipette tip.

The single-molecule fluorescence signal was distinguished from the background at the pipette tip based on characteristic single-molecule spectroscopic signatures: fluorescence blinking, single-step photobleaching, specific polarization orientation, and, most importantly, simultaneous single-channel ion current recording. The last simultaneous verification was that only a single-channel ion current associated with a single ion channel in the patch was observed during fluorescence imaging. The signal-to-noise ratio was typically within a range of 5–10 in the control LB bilayer imaging at the LB bilayer or PMMA polymer film, and 3–6 in the PCFM imaging at the pipette tips.

RESULTS AND DISCUSSION

Single-molecule spectroscopy is powerful in resolving complex and inhomogeneous dynamics and kinetics of protein systems. However, the fluctuation of most of the measured parameters is intrinsically pertinent to single-molecule measurements (Zwanzig, 1990; Lu and Xie, 1997; Xie and Trautman, 1998; Moerner, 2002). For example, single-molecule fluorescence has spectral and intensity

fluctuations, FRET efficiency fluctuations, and lifetime fluctuations. These fluctuations are not observable in a typical ensemble-averaged experiment, and in most cases they are associated with single-molecule conformations and local environments. A reliable conclusion should typically be drawn from sound statistical analyses of a sufficient number of measurements of single-molecule events. Although the fluctuations give a larger standard deviation in the data of a single-molecule measurement compared to that of an ensemble-averaged measurement, the information obtained from single-molecule experiments has not been obtainable from ensemble-averaged experiments.

Heterodimers of gramicidin statistically correlate with maximum spFRET at an open state

Using PCFM, we conducted colocalization and spFRET imaging experiments simultaneously with single-channel patch recording. Asymmetric incorporation of Cy5-gramicidin C and TMR-gramicidin C led to the formation of single channels tagged with two different dyes, the donor and acceptor of the fluorescence energy transfer. Colocalization and spFRET within a single gramicidin heterodimer were probed by dual-color excitation and donor-acceptor two-channel fluorescence imaging, using an instrument configuration similar to one reported previously (Cognet et al., 2000). Fig. 3 *a* shows two fluorescence images taken at the tip with 514-nm (image 1) and 632-nm (image 2) excitation wavelengths, using a filter that allows imaging of both Cy5 and TMR emissions, both of which are clearly discernable in Fig. 3 *a*. From the 2-D Gaussian fitting of the colocalized signals, the two dye molecules were found to be within the fluorescence imaging diffraction-limited spot (~ 300 -nm width), which suggests that a pair of colocalized TMR-gramicidin and Cy5-gramicidin monomers was present at the tip when the active single-channel current signal was observed (Fig. 3 *a*). The colocalization imaging measurement is important and was used periodically to check the coexistence of the donor and acceptor molecules and to differentiate the situations of non-FRET from acceptor photobleaching events. To further characterize the heterodimerization states of TMR-gramicidin and Cy5-gramicidin, spFRET images were collected using a 645-nm long-pass filter, allowing only the fluorescence of acceptor Cy5 to pass. Toggling between 514-nm and 632-nm excitation wavelengths enabled the measurement of spFRET and verification of the coexistence of the single acceptor (Fig. 3 *b*). The spFRET that occurred during the open states of the channel is shown in Fig. 3 *b*. By counting images correlated with the electric signal that reflected the open-closed activity of single channels at the patch, we found that $\sim 70\%$ of channel-opening events were correlated with spFRET. Estimating that $\sim 25\%$ of the molecules undergo blinking events, we attributed the small fraction of non-FRET to acceptor photobleaching. Interestingly, we also observed that spFRET

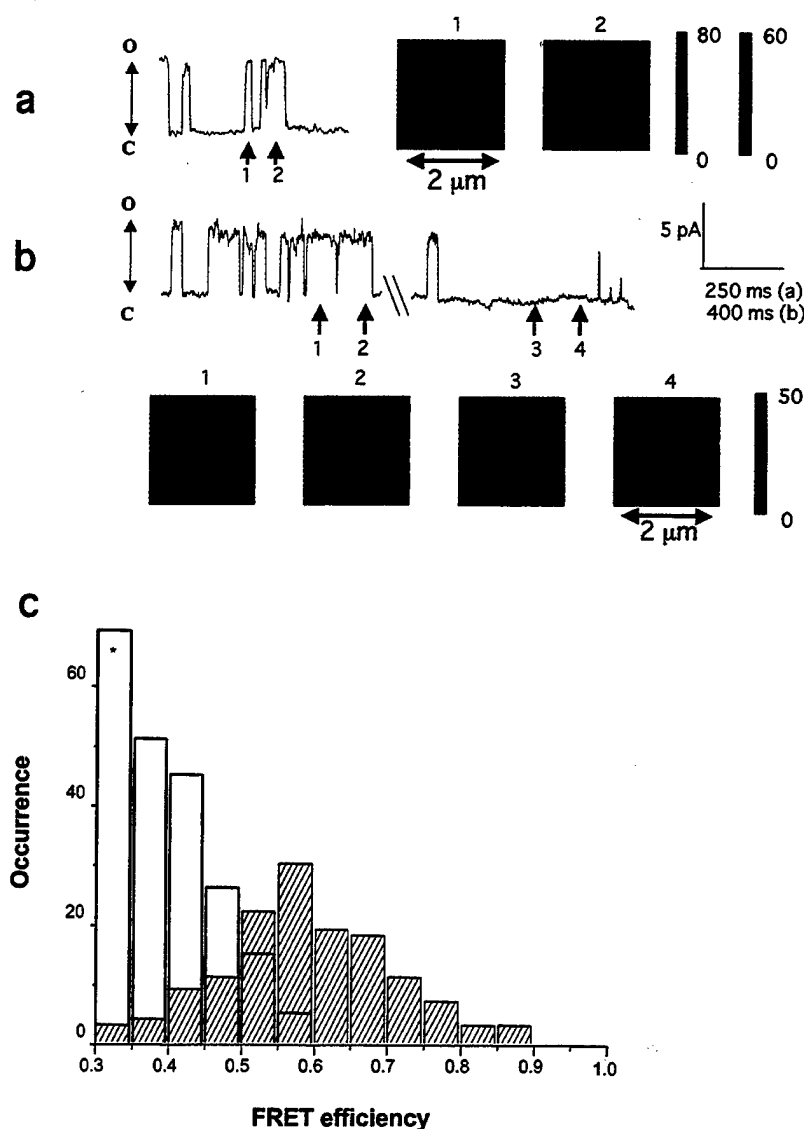


FIGURE 3 PCFM measurement of single TMR/Cy5-gramicidin heterodimer ion channels. Green and red arrows mark the timing of the toggled excitation at 514 nm and 632 nm, respectively. (a) Colocalization of two gramicidin monomers labeled with two different dyes, TMR and Cy5. Single-channel current trajectory recorded at 100-mV polarization potential, with arrows indicating the point of exposure of two consecutively recorded frames. The first frame was taken at 514-nm excitation (with the OG550 long-pass filter), indicating single-molecule fluorescence of TMR. The second frame was taken at 632-nm excitation, indicating single-molecule fluorescence of Cy5. From a 2-D Gaussian fitting, the molecules were found to be within 30 nm of each other (the minimal positional accuracy allowed by the measurement system). "C" and "O" denote the current corresponding to closed and open states, respectively. (b) spFRET within a pair of gramicidin monomers labeled with two different dyes, TMR and Cy5. Single-channel current trajectory (upper panel) with four arrows indicating the exposure of four consecutively acquired frames (lower panel). Frames 1 and 3 were taken at 514-nm excitation with a red-emission long-pass filter (RG645). Frames 2 and 4 were taken at 632-nm excitation with a red-emission long-pass filter (RG645). Frame 1 indicates that FRET occurred between the TMR/Cy5 spFRET pair when the gramicidin channel was at an open state. Frame 2 verifies that a single Cy5 molecule was present. Frame 3 indicates that no spFRET occurred when the channel was at a closed state. Frame 4 verifies that the Cy5 was still present. Subsequent images taken without the red long-pass filter (RG645) did show colocalization as shown in a. (c) FRET efficiency distribution of "channel open" TMR/Cy5 heterodimers (shaded bars) with mean of 0.59, width of 0.21, and "channel closed" (open bars) with a maximum value of 0.57. An asterisk is inserted to indicate the 70% of channels showing no appreciable FRET (below 0.3) in the channel-closed state (or 30% of channels in the channel-open state).

occurred in $> \sim 10\%$ of the cases where the channels were closed. We have further verified the FRET anticorrelation of the donor-acceptor fluorescence intensities by photon time-stamping, time-correlated single-photon counting (Hu and Lu, 2003) and proved the presence of FRET (Harms et al., 2003; G. S. Harms, G. Orr and H. P. Lu, unpublished results).

A broad distribution of FRET efficiency (Ha et al., 1999; Brasselet et al., 2000; Cognet et al., 2000),

$$F = \left(1 + \frac{\eta_{\text{Cy5}} \Phi_{\text{Cy5}} I_{\text{TMR}}}{\eta_{\text{TMR}} \Phi_{\text{TMR}} I_{\text{Cy5}}} \right)^{-1},$$

was found at gramicidin heterodimers, and the efficiency was evaluated by the activity of the channels at the time that the images were taken (Fig. 3 c). I_{TMR} and I_{Cy5} are the total photon counts for single-molecule images of TMR and Cy5,

η is detection efficiency, and Φ is the quantum yield. We used the values of η_{TMR} and η_{Cy5} discussed above as well as $\Phi_{\text{TMR}} = \Phi_{\text{Cy5}} = 0.28$ (Cognet et al., 2000). When the channels were open, the Gaussian-like distribution of the efficiency showed a mean value of 0.59 ± 0.05 with a full-width half-maximum (FWHM) of 0.21. The standard deviation of the F was estimated to be ± 0.05 based on shot-noise error propagation analysis (Press et al., 1990). Our control experiment of polarization-dependent imaging (Harms et al., 2003) indicated that the rotational diffusion of the TMR and Cy5 dye molecules had a much faster rate than the imaging rate, so that the orientation effect of the dye dipoles on the FRET efficiency was averaged out during the 5-ms imaging collection time for each frame. The mean spFRET efficiency and the FWHM of 0.21 correspond to a mean donor-acceptor distance (d) of $\sim 56 \text{ \AA}$ and a range of $\sim 52 \text{ \AA} \leq d \leq \sim 63 \text{ \AA}$, respectively. However, $\sim 30\%$ of

the images obtained from the channel open states showed no detectable FRET efficiency, mostly because of photobleaching and spectral fluctuation. We cannot differentiate the data between no FRET and low-efficiency FRET below 0.3, due to the experimental limitation of the signal-to-noise ratio of the single-molecule PCFM imaging. The distribution of FRET efficiency of the closed state of single channels had a mean of 0.42 ± 0.05 (excluding the low signal data) and extended to a maximum efficiency of 0.57. Therefore, the difference in the FRET between open and closed states was statistically significant and beyond the measurement noise. The broad FRET distributions of both open and closed states of the single gramicidin channels suggest that the channels could be in closed states even when the two gramicidin monomers are still in an intermediate dimerized state (Fig. 4, *a* and *b*). We postulate that the partially dissociated dimer may have had a distorted ion channel pore, creating a significant change in the channel electric current, though the physical distance between the two monomers did not significantly change (Fig. 4 *b*). It has been suggested that

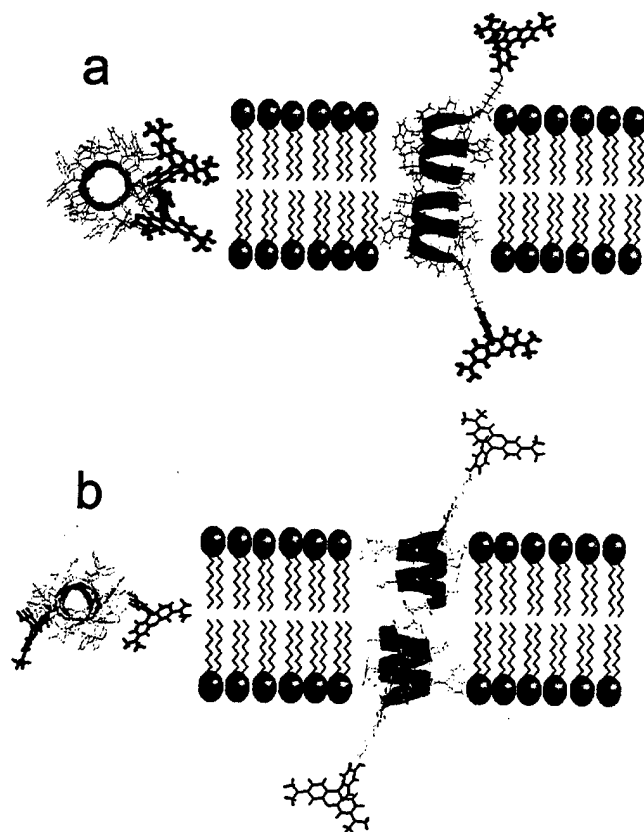


FIGURE 4 Molecular structures (top and side views) of the possible open (*a*) and closed (*b*) conformational states of a gramicidin dimer in a lipid bilayer. The molecular structures were calculated by molecular dynamics simulation using Discovery and Insight II software. It is evident that slight rotational distortion along the dihedral angle can significantly narrow the channel pore size and separate the dye probe pair.

there are six hydrogen bonds at the N-terminal head-to-head cross-linking the dimer (Szabo and Urry, 1978; Koeppe and Andersen, 1996). It is possible that one or more broken hydrogen bonds can give different conformations of the pore structure of a gramicidin dimer. In Fig. 4, we show that the energy-minimized structure of gramicidin heterodimer FRET pair (represented by using TMR molecular structures) has the minimum donor-acceptor distance of ~ 50 – 52 Å. The relative dihedral angle between the two monomers at different conformations of the conducting dimer (Fig. 4) can give a distance change up to 6 Å, which can result in a 0.15 change in FRET efficiency. We note that the calculated spFRET efficiency is only reliable in the range of $0.3 < F < 0.7$ due to the measurement noise associated with the effects of shot noise and photophysical dark states. Nevertheless, spFRET imaging results (Fig. 3 *c*) provide unambiguous evidence that the gramicidin channel dynamics involve multiple open and closed states associated with different conformations of the gramicidin dimers.

The spFRET imaging measurement using toggled dual-color excitation and simultaneous correlation with the single gramicidin ion channel patch electric recording is a novel approach, and the results of the correlation between the spFRET and the open-closed states of the heterodimer gramicidin strongly suggest the existence of more than one conformational state at each open and closed state of the gramicidin ion channels in the lipid bilayer. These hidden multiple conformational states are not observable from ensemble-averaged measurements, and also are "silent" and not resolvable in a conventional patch recording analysis.

Homodimers of dye-labeled gramicidin statistically coincide with maximal fluorescence self-quenching at an open state

We have obtained fluorescence images of single TMR-gramicidin homodimers at bilayer patches simultaneously with measurements of their single-channel electric currents using PCFM. A segment of a single-channel electric current trajectory (Fig. 5 *a*) with synchronous fluorescence images (Fig. 5 *b*) illustrates the correlation between the transitions of the channel from the closed to the open state and fluorescence hot spots (Fig. 5). Single hot spots were analyzed for width (Fig. 6 *a*) and intensities (Fig. 6, *b* and *c*). These mean values are from single homodimers of the dye-labeled gramicidin, an interpretation based on the coincidence of the recorded single-channel electric currents with the fluorescence images and on the various control experimental analyses discussed above.

We further observed that the width of the hot spot fluorescence increased with fluorescence intensity (Fig. 6 *a*). Theoretically, the spread functions of the one-point-source and two-point-source images are Gaussian ($G(x,y) = A \exp[-(x-x_0)^2 + (y-y_0)^2]/B$) and a convolution of two

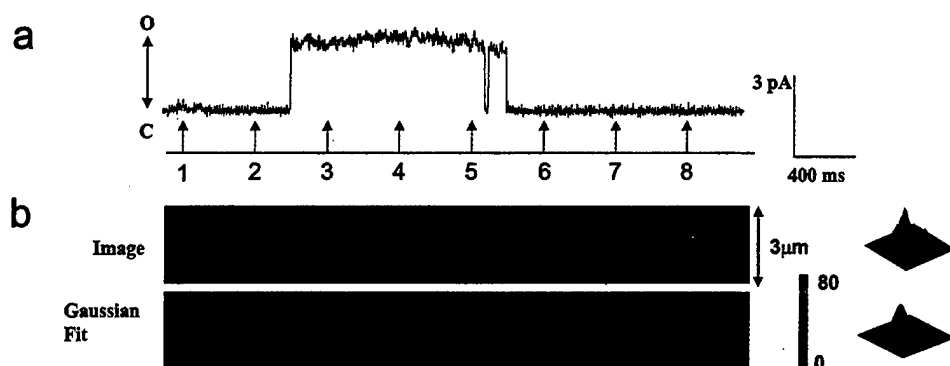


FIGURE 5 PCFM measurement of single TMR-gramicidin homodimer ion channels. (a) Upper trace: single-channel electric current trajectory recorded at 50-mV polarization potential from a single TMR-gramicidin channel at a patch-clamp pipette tip. Lower trace: time flags indicating the 5 ms of laser excitation and CCD camera exposure simultaneously acquired with the single-channel electric recording. "C" and "O" denote the electric current corresponding to closed and open states, respectively, of the single channel.

(b) Top panel: Consecutive fluorescence images of the pipette tip taken simultaneously with the single-channel electric current recording shown in *a*. Bottom panel: plot of the 2-D Gaussian fitting for the above images. Right inset: 3-D plot of the first image (*top*) and 3-D plot of the 2-D Gaussian fitting (*bottom*) with subtracted background intensity added to the base shown in gray.

Gaussian functions ($G \times G$), respectively. Assuming that the imaging spatial resolution is optical diffraction-limited, the FWHM is a constant, $\sim 2B^{0.5}$, for $G(x, y)$ at different signal intensity and the FWHM of $G \times G$ is dependent only on the two-point-source separation. The error bars for data points in Fig. 6 *a* are contributed from the shot noise, finite signal-to-noise ratio, digitized photon counting, and digitized pixels. The widths of the hot spots were essentially diffraction-limited but wider when the channels were fully closed (Figs. 5 *b* and 6 *a*). This observed behavior is beyond the standard deviation (Fig. 6 *a*). These observations are consistent with the expectation that the channel is fully open when the dimer is fully associated, leading to maximal fluorescence self-quenching because the two labeled dye molecules have the closest distance across the lipid bilayer at the patch (Fig. 4 *a*). Rarely (<5% of images) were two separate and spatially resolved imaging spots observed when a single channel was closed, i.e., a dissociated dimer. In most cases, the two dissociated monomers do not separate beyond the imaging spatial resolution, ~ 300 nm, or even beyond the diffraction limit. The imaging signal-to-noise ratio is not high enough for obtaining a clear image of two separate spots, as we observe that the widths of the spots are larger than the diffraction limit at higher fluorescence intensity (Fig. 6 *a*).

We found that the diffusional motion of single gramicidin channels at the dPhPE:DPhPC bilayers was confined within the imaging spatial resolution. According to viscosity studies, dPhPE is likely to be in a gel-phase at room temperature (Janko and Benz, 1977). Based on our control measurements in a large, supported LB-membrane bilayer, the diffusion coefficient of TMR-gramicidin was estimated to be $<10^{-10}$ cm²/s in a dPhPE LB layer, yielding ~ 0.2 - μ m average displacement in 1 s. This is consistent with the fact that we rarely observed diffusion of a dissociated dimer into two resolved monomers. Although the diffusion coefficient of the TMR-gramicidin can be as high as 10^{-8} cm²/s in DPhPC at room temperature (Borisenko et al., 2003), the 4:1 dPhPE:DPhPC bilayers may have small domains (<500 nm)

of the DPhPC formed (Heller et al., 1998), confining the diffusional motions of the TMR-gramicidin molecules within the DPhPC. Such "corrallled" motion may be accountable for the rare observation of an optically resolved separation of a pair of TMR-gramicidin monomers dissociated from a dimer. It has been known (Heller et al., 1998) that domains of DPhPC can form in DPhPE bilayers and that gramicidin channels typically prefer to stay in the DPhPC regions.

The diffraction-limited single hot-spot images of TMR-gramicidin dimers also reflected different degrees of fluorescence self-quenching (Zhuang et al., 2000). The self-quenching critical radius of the rhodamine 6G dyes is ~ 46 Å (Penzkofer and Lu, 1986). The fluorescence self-quenching between the two dye molecules causes an intensity decrease as the two monomers become closer and form a conductive channel. The average distance between the transition dipoles of the TMR dyes was calculated to be ~ 50 Å, and the transition dipoles were nearly aligned in the fully dimerized state (Fig. 4 *a*).

The mechanism of fluorescence self-quenching is still largely unknown and system-dependent (Zhuang et al., 2000; Penzkofer and Lu, 1986). In our gramicidin ion channel system, because the two TMR dye molecules are not in contact to form a dimer or excimer, the possible mechanism of the observed weak self-quenching effect is likely associated with reabsorption, and secondary fluorescence depends on the fluctuating orientation of the two TMR dye molecules across the lipid bilayer.

Fig. 6, *b* and *c*, shows the distributions of the fluorescence imaging intensity in correlation with the open and closed states of the gramicidin channels, measured by PCFM. The intensity of the hot spots was statistically higher when the channel was closed (Fig. 6 *b*), and lower when the channel was open (Fig. 6 *c*). The means of the distribution were found to be 67 ± 8 counts/ms at closed states and 52 ± 7 counts/ms at open states. Furthermore, the difference in fluorescence intensities of the single-channel images taken

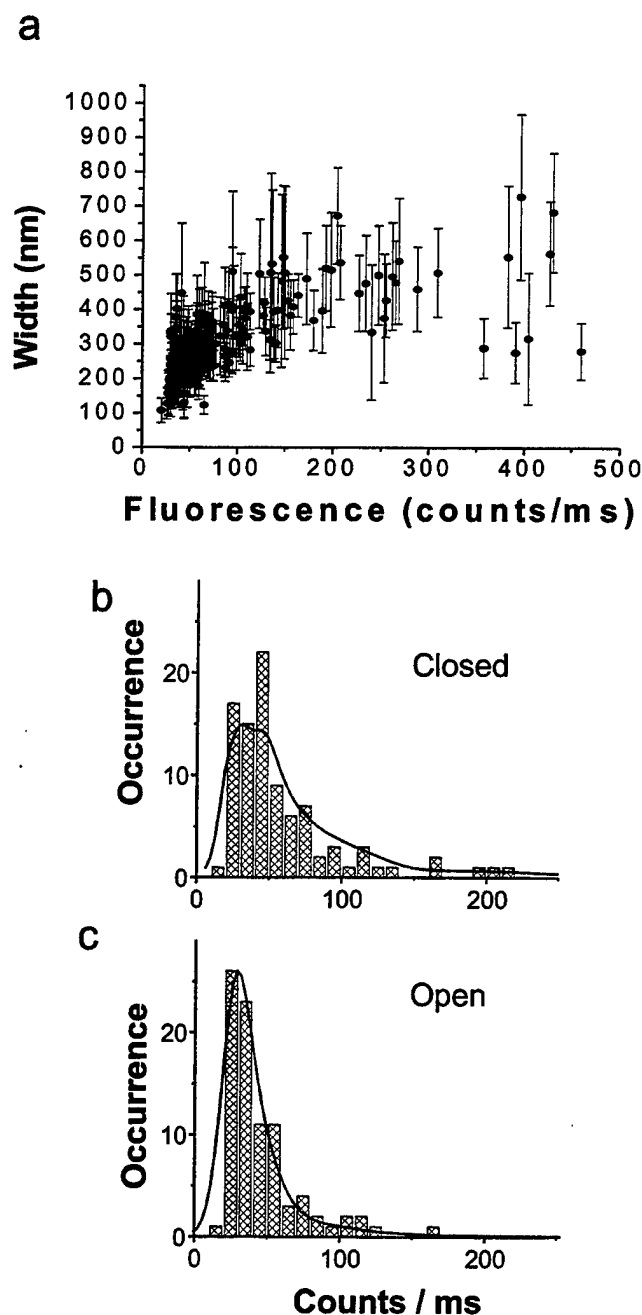


FIGURE 6 (a) Fluorescence intensities of single hot spots versus fluorescence imaging width of the spots as derived from 2-D Gaussian fit. (b, c) Distributions of the single-channel fluorescence intensity for closed and open states of single channels with PDF and fitting.

for the open and closed states was also found to be statistically significant, based on the paired *t*-test ($t = 2.58$, $P = 0.0297$). We also noted that the broadness of the distribution is significant and beyond the measurement noise and that both quenched and unquenched gramicidin channel dimers probably existed in the channel closed state. Because the efficiency of self-quenching depends on the distance

and the relative orientation between the two molecules, the broader distributions of the fluorescence intensity at the closed (Fig. 6 *b*) and open (Fig. 6 *c*) states also suggests that the two monomers form "closed" or "open" dimers of different conformations (Fig. 4). This result is consistent with the spFRET imaging result (Fig. 3), also suggesting that there are multiple intermediate conformation states corresponding to different distances between the homo-pair of TMR molecules in a single gramicidin channel based on the wide distribution of the fluorescence self-quenching efficiency among the single gramicidin channels in the lipid bilayer.

Autocorrelation analysis of single-channel electric current trajectories characterizes the dynamics of conformational changes of the multiple closed and open states

To further explore the possible existence of the multiple conformational states involved in gramicidin ion channel dynamics, we applied autocorrelation function analysis (Zwanzig, 1990; Neher and Stevens, 1977; Labarca et al., 1985; Lu et al., 1998) of the single-channel electric current trajectories. Detailed single-channel current analyses support our attribution that multiple conformational states are involved in the spatially confined gramicidin dimer channel dynamics. Fig. 7 *a* shows a typical single gramicidin channel electric current trajectory. The intermediate conductive states, in addition to the fully open and closed states, are evident in the trajectory and are further revealed in an electric current amplitude distribution shown in Fig. 7 *b*. There is a significant population beyond the measurement noise and within the time resolution in the distribution between the typical open-closed electric current amplitude bimodal distribution (Fig. 7 *b*). The autocorrelation function deduced from the single-channel electric current trajectory (Fig. 7 *a*) shows nonexponential decay (Fig. 7 *c*), and the nonexponential autocorrelation functions were commonly observed for both dye-labeled and non-dye-labeled gramicidin channels (D. J. Panther, G. S. Harms, G. Orr, and H. P. Lu, unpublished results). The nonexponential autocorrelation functions suggest that the transitions between multiple open-closed states of a single channel either occur at a fluctuating rate (Zwanzig, 1990; Xie, 2002; Yang and Xie, 2002; Jung et al., 2002) or involve a complex kinetic mechanism associated with multiple intermediate states (Schenter et al., 1999; Jung et al., 2002; Reilly and Skinner, 1993). We have analyzed the dynamics using an autocorrelation function of sequential open dwell-time (index number, m) $\{t_{on}(m)\}$, calculated by $\langle \Delta t_{on}(0) \Delta t_{on}(m) \rangle$, where $\Delta t_{on}(m) = t_{on}(m) - \langle t_{on}(m) \rangle$ (Fig. 7 *d*). The single-exponential decays that were fitted to the autocorrelation functions of $t_{on}(m)$ (Fig. 7 *d*) and $t_{off}(m)$ (not shown) were $2.6(\pm 1.0) \times 10^{-2} \text{ ms}^{-1}$ and $0.5 \pm 0.2 \text{ ms}^{-1}$, respectively. The open and closed dwell-time

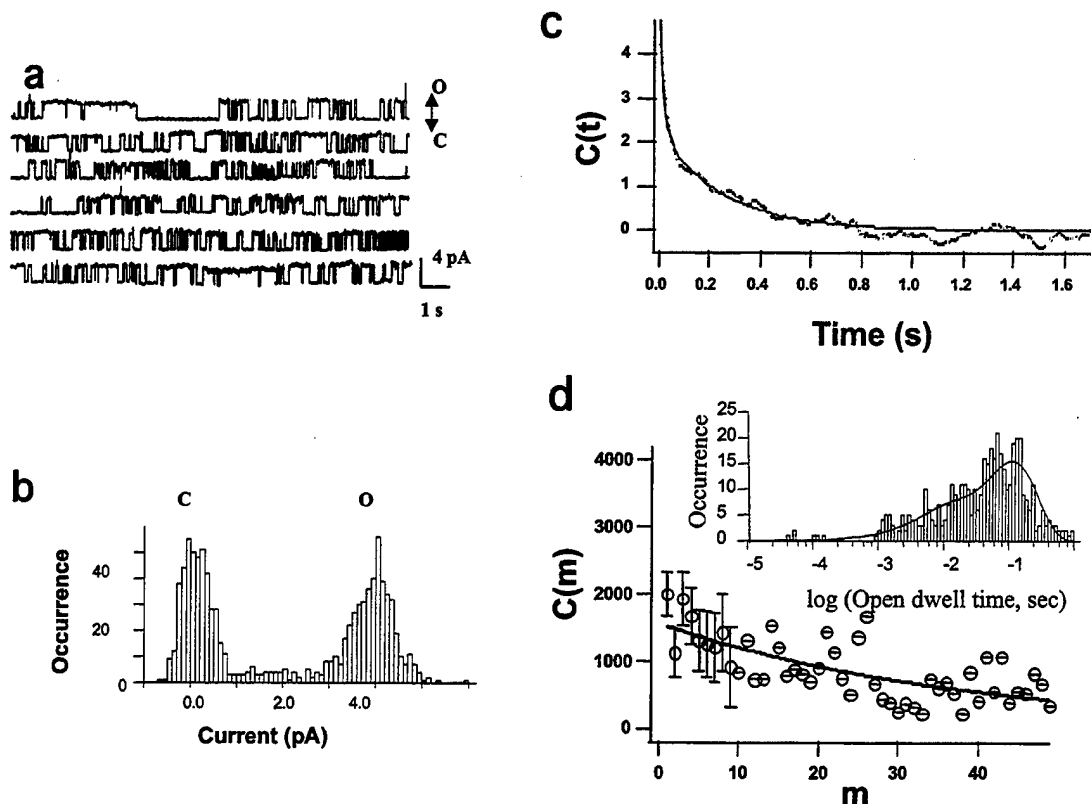


FIGURE 7 (a) Single-channel electric current trajectory, $\{i(t)\}$, recorded at 100-mV polarization potential from a single gramicidin channel in a lipid bilayer patch (4:1 dPhPE:DPhPC) formed at a patch-clamp pipette tip. The single-channel electric current trajectory indicates that there are only single-channel events (O'Connell et al., 1990). (b) Electric current amplitude histogram calculated from the single-channel trajectory in *a*. Intermediate electric current amplitude reflecting the inhomogeneous conductivity is evident although the channel shows essentially two-state kinetics. "C" and "O" denote the current corresponding to closed and open states, respectively. (c) Autocorrelation function, $C(t) = \langle \Delta i(0)\Delta i(t) \rangle$, calculated from the single-channel trajectory in *a*. Biexponential fitting gives $k_1 = 4.0 \pm 1.0 \text{ s}^{-1}$, $A_1 = 0.48$, and $k_2 = 53 \pm 5 \text{ s}^{-1}$, $A_2 = 0.52$; using $A_1 \exp(-k_1 t) + A_2 \exp(-k_2 t)$. (d) Autocorrelation function of "open" dwell times, $r(m) = \langle \Delta i(0)\Delta i(m) \rangle$, where m is the index of the "open" dwell-time trajectory for the single TMR-gramicidin ion channel recorded in *a*. The single-exponential decay rate constant is $2.6 (\pm 1.0) \times 10^{-2} \text{ m}^{-1}$, and the first point (not shown) of the autocorrelation contains the uncorrelated noise and fast fluctuation beyond the instrument resolution. Error bars are shown for the first ten data points. The calculated autocorrelation times are in the timescale of seconds and subseconds (*c* and *d*). The channel open-closed dwell times are also at subsecond timescale (*a*). Using a 4-kHz filter instead of a 6-kHz filter does not affect the results, even if the possible aliasing effect exists at millisecond timescale. Inset: The distribution of "open" dwell time deduced from the same trajectory in *a*. The solid line shows a distribution envelope fitting with a bi-Gaussian function with a relative weight of 0.808 and 0.192 at averaged dwell time of 1.13×10^{-1} and $8.67 \times 10^{-3} \text{ s}$, respectively. Inhomogeneous distribution is evident.

correlations suggest that the channel open and closed states are dependent within a short period of "memory" time (Lu et al., 1998; Zwanzig, 1990; Yang and Cao, 2001; Vlad et al., 2002; Jung et al., 2002; Xie, 2002; Yang and Xie, 2002). The memory time for the open dwell-time was determined to be $\sim 8 \text{ s}$ (38 cycles), but only $\sim 0.4 \text{ s}$ (8 cycles) for the closed dwell-time. We postulate that the "memory" effect originates from the channel open-closed conformational motions that are trapped in metastable processes under different rates of toggling among different conformational states. This dynamically inhomogeneous behavior, also defined as dynamic disorder (Zwanzig, 1990), has been observed in other ion channel systems (Sakmann and Neher, 1995; Neher and Stevens, 1977). Recently, single-molecule spectroscopy has made progress in understanding this dynamic disorder

behavior, which is beyond the scope of the conventional kinetics (Lu et al., 1998; Zwanzig, 1990; Yang and Cao, 2001; Vlad et al., 2002; Jung et al., 2002; Xie, 2002; Yang and Xie, 2002). Typically, the mechanism of the dynamic disorder can be analyzed by the "Agmon-Hopfield" diffusive model (Agmon and Hopfield, 1983; Agmon, 2000) or by the multiple-state Markovian models (Schenter et al., 1999; Sakmann and Neher, 1995; Neher and Stevens, 1977). The Fig. 7 *d* inset shows the distribution of the "open" dwell-time. At least two subgroups of "open" dwell times contributed to this broad distribution. We postulate that the dynamic disorder of this gramicidin system in dPhPE:DPhPC bilayers can be characterized by a simple coupled Markovian kinetic model, for example, the 2×2 model (Schenter et al., 1999), which describes two slowly

interconverting open-closed kinetic processes that have different kinetic rates.

The open and closed dwell times are found to be dependent over periods of hundreds of milliseconds, which suggests a trapped or not fully dissociated subset of channel conformers. We note that the gramicidin channel dynamics can be significantly different using different lipid bilayers because the channel open and closed activities are associated with the gramicidin diffusion motion in a particular bilayer. Nevertheless, the gramicidin ion channel dynamics at the dPhPE:DPhPC bilayers are apparently associated with more than two states: 1), The open states of the gramicidin dimer involve slow, strongly correlated conformational fluctuations that lead to dynamic disorder of the dissociation rate (Lu et al., 1998; Zwanzig, 1990; Yang and Cao, 2001; Vlad et al., 2002). 2), Gramicidin dimerization involves multiple, weakly correlated or noncorrelated association kinetics at different rates. Although the association kinetics of gramicidin have been studied and discussed in terms of fast fluctuations (Sigworth et al., 1987) and the existence of possible "mini-channels" (Busath and Szabo, 1981; Sawyer et al., 1989), our observations of the "memory" effect in the gramicidin channel dynamics in dPhPE:DPhPC bilayers provide a new insight into the molecular level. However, we are not sure, at this stage, if the complex mechanism we observe is generally true for the gramicidin ion channels in other lipid bilayer systems.

A multistate model for gramicidin channel dynamics

Based on the information derived from the PCFM imaging, we propose a working model of multi-state gramicidin channel dynamics at dPhPE:DPhPC bilayers (Fig. 8). The open (*left*), closed (*right*), and intermediate (*middle*) states can be presumably the outcome of intra- and intermolecular conformational changes, geminate recombination, and non-geminate recombination. Although the valine (Townsend et al., 2001; Sigworth et al., 1987) and tryptophan residues (Townsend et al., 2001) may undergo large structural fluctuations and alter channel conductance by perturbing the structure of the dimeric channel (Koeppel and Andersen, 1996; Townsend et al., 2001; Koeppel et al., 2000; Salom et al., 1995; Woolf and Roux, 1994), we postulate that the intermediate conformers result predominately from the six intermolecular H-bond fluctuations that disrupt the pore structure of the gramicidin dimer. Conformational changes at the N-terminal linkage of the dimer may significantly disrupt the column of structured water, the channel cavity structure, and the electrostatic field, thereby perturbing channel conductance (Eisenberg, 1998). Drastic conductance modifications were reported for gramicidin channels with mutated N-terminal head-to-head connections (Sigworth et al., 1987; Szabo and Urry, 1978), indicating that the six intermolecular H-bonds formed at the contact region of the dimer stabilize the fully conductive gramicidin channel

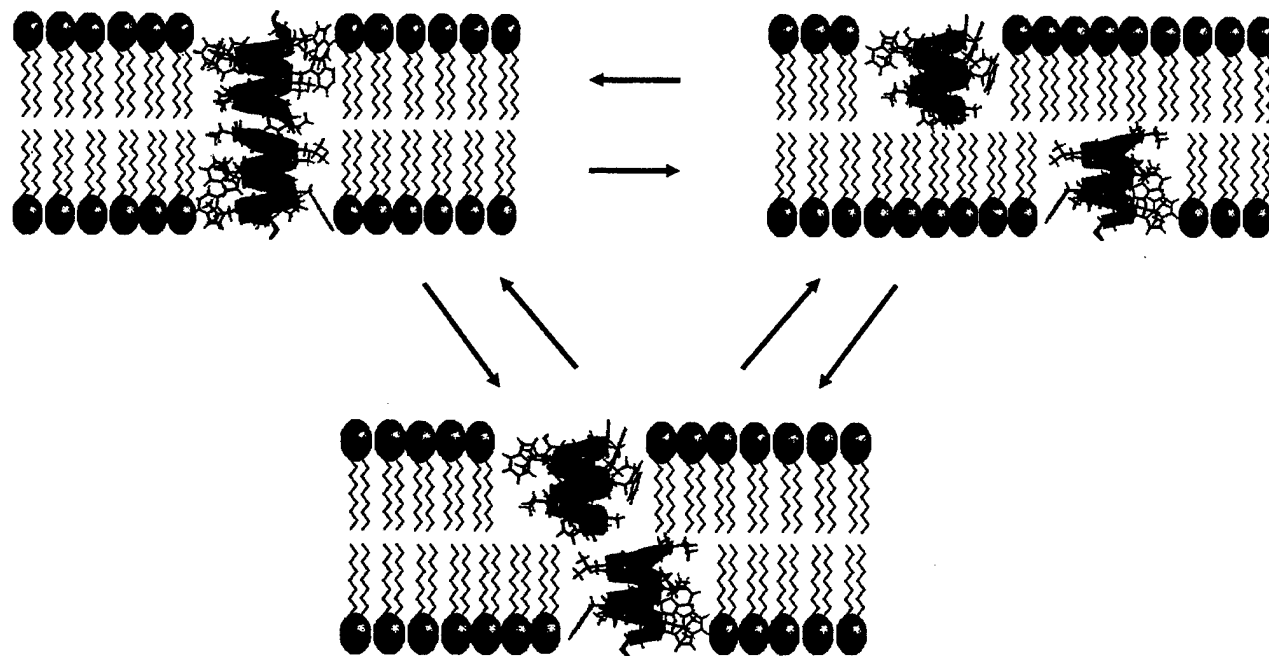


FIGURE 8 Proposed multistate model of gramicidin dynamics indicating a third "multistate" of nonconducting or partially conducting channels with larger separation between the fluorophore probes. In this model, the third "multistate" represents more than one intermediate conformation state. Alternative rotationally distorted conformations are possible for the fully conductive state of the gramicidin channels (Roux and Karplus, 1994).

(Szabo and Urry, 1978; Koeppe and Andersen, 1996). Partial disruption of these H-bonds results in incomplete association or dissociation of the dimers (Szabo and Urry, 1978). Given that each H-bond has $\sim 3\text{--}5$ kcal/mol dissociation activation energy, the dissociation of these bonds plausibly occurs in the millisecond timescale at room temperature. These estimated H-bond dissociation times are in the same timescale as the gramicidin dimer changes between open and closed states. The conformational changes that could originate from intramolecular changes or lipid-channel interactions are expected to occur at a faster timescale and, therefore, should not result in the observed self-quenching and FRET effects at the millisecond timescale. Therefore, we hypothesize that the conformational changes that we measured arise from the N-terminal H-bond dynamic changes. We are not able to provide direct structural evidence of the H-bond fluctuation using our current PCFM. The structural and dynamical information may be obtained by integrating a Raman spectroscopic probe with the PCFM, which is beyond the scope of this article.

Diffusion of individual monomers that dissociate from the dimerized state (Tank et al., 1982) is confined within the optical diffraction-limited spot size, based on our experimental observation using wide-field imaging. This observation has been further confirmed by time-dependent and static anisotropy analysis of gramicidin C-TMR in the LB bilayer at the patch-clamp tips. The detailed results and discussion will be reported elsewhere. Briefly, the decay time of the anisotropy is ~ 5 ns, and the static anisotropy is 0.25 ± 0.05 (Harms et al., 2003; G. S. Harms, G. Orr, and H. P. Lu, unpublished data). Based on our observation, in most of the events, the pairs of gramicidin monomers did not diffuse away from each other beyond the imaging diffraction-limited scale (≤ 300 nm), and we postulate that the geminate recombination may be associated with single-channel conformational changes. It is possible that the diffusion of gramicidin monomers was confined to domains (≤ 300 nm) likely associated with the inhomogeneity (Nielsen and Andersen, 2000) of the dPhPE:DPhPC bilayers, as discussed above.

CONCLUSIONS

We have developed a new single-molecule approach, PCFM, combining single-molecule fluorescence imaging microscopy and single-channel patch recording to simultaneously record the single-channel trajectories of fluorescence imaging and channel electric current recording. We have demonstrated the application of PCFM in studying the ion channel dynamics of single gramicidin dimers in a lipid bilayer at the patch-clamp tip. By correlating single-channel fluorescence quenching and spFRET imaging with single-channel electric current recording simultaneously acquired, our results strongly suggest the existence of multiple conformational states of a single gramicidin channel associated with the channel open-closed activity at a dPhPE:DPhPC

bilayer. Although the inhomogeneous dynamics of the conformational change were probed in a lipid bilayer, the exact structures of the conformational states require further studies, for example, by correlated Raman spectroscopy. The new single-molecule experimental approach for studying ion channel conformational dynamics and mechanisms should be applicable for direct study of ion channel proteins in living cells (Orr et al., 2002).

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